

*Full Length Research Paper*

# Genetic analysis of twenty two selected genotypes of *Jatropha curcas* L. (physic nut) from Africa, Asia and America, using SSR and AFLP markers

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Twenty two genotypes of *J. curcas* L. from Africa (Senegal, Burkina Faso, Mali, Congo and Madagascar Island), Asia (Cambodia, China and India) and America (Ecuador, Dominican Republic and Brazil) selected for their vigor and their productivity were analyzed with ten SSR primer pairs and six AFLP primer combinations. The two marker approaches showed their ability to effectively reveal polymorphism among the selected genotypes: 94.02 and 56% of polymorphism for AFLPs and SSRs respectively. Among the three groups of selected genotypes, the Asian group was the least diverse while the genetic diversities found in African and American groups were slightly comparable. The Nei's genetic diversity ( $H_e$ ) of all twenty-two selected genotypes was 0.2029 based on combined SSR+AFLP data. The  $G_{st}$  value and the AMOVA analysis indicated that more than 80% of the genetic diversity resided within the groups. The analysis of the genetic relationships between the genotypes using the Nei's standard dissimilarity matrix gave dissimilarity coefficients ranging from 0.14397 to 0.73943 with an average of 0.3540. The most distant genotypes were found between a genotype from Africa (Congo) and one from America (Ecuador). The clustering of genotypes obtained with the neighbor-joining dendrogram and the PCoA analysis revealed the existence of a certain level of diversity that can be used by breeders.

**Key words:** Biodiesel, genetic diversity, jatropha, molecular markers, AFLP, SSR, plant breeding.

## INTRODUCTION

*Jatropha curcas* L. is a small perennial tree native to tropical America that has spread to tropical and subtropical regions of Asia and Africa over the last 200

years (Marinho et al., 2018). International attention to this plant has emerged during the past decade from the need to reduce dependence on the increasing scarcity and

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cost of fossil fuels and their associated carbon emissions (Moniruzzaman et al., 2017). *Jatropha* has been identified as a promising candidate for clean and renewable energy because of its seed oil which had the characteristics of diesel, its non-food nature and its supposed capability of high yield on marginal soils with minimum management requirements and inputs (Vásquez-Mayorga et al., 2017; Cardoso et al., 2018). For these reasons, *jatropha* was assumed to have potential to bring socio-economic benefits and there has therefore been a heightened global interest in large-scale *jatropha* cultivation and many *jatropha* projects have been implemented in Asia, Africa and Latin America (Soto et al., 2018). However, the performance of this new crop has been very disappointing despite claims of high yields, drought tolerance, low nutrient requirements, high returns on investment etc.; *jatropha* investments failed to meet global expectations (Antwi-Bediako et al., 2019). Consequently, doubts have been cast on the profitability and the financial viability of its cultivation, leading to the disillusionment of farmers and disadoption of *jatropha* by most of them (Soto et al., 2018; Vandepitte et al., 2019).

One of the main causes of the problem of *jatropha* cultivation was the low seed yield (Moniruzzaman et al., 2017; Soto et al., 2018). The cultivars used did not achieve the initially anticipated high seed yields, because the expectations were not based on field performance and validated scientific knowledge (Govender et al., 2018). The promotion was based on a multitude of unfounded claims, which did not lead to a commercially successful *jatropha* production.

Actually, *J. curcas* is an incipiently domesticated species with no availability of competitive commercial cultivars (Díaz et al., 2017). It is mostly a xenogamic plant, highly heterozygous for most characteristics, which implies a high degree of segregation (Díaz et al., 2017). The less productivity observed was because of unavailability of suitable high yielding varieties and a good commercial variety is still missing (Cardoso et al., 2018). As for all non-domesticated perennials, time is needed for the development of commercial varieties of *jatropha*; unfortunately this fact seems to have been overlooked and *jatropha* has been considered a commercial crop without any real scientific basis.

According to Tjeuw (2017), *Jatropha* has the potential to be a cash-crop like cotton or rubber (*Hevea brasiliensis* M.) which are both equally non edible but seem to be well accepted and not seen as a threat to food production. However for the successful exploitation of *jatropha*, there is an urgent need for improved genetic material with known yield characteristics over a wide range of climatic and agro-ecological conditions. It is one of the main effective solutions and the future challenge for *jatropha* breeders (Díaz et al., 2017; Moniruzzaman et al., 2017; Cardoso et al., 2018). This will require extensive

plant breeding and molecular strategies to investigate systematically and scientifically *jatropha* potential for the development of a competitive commercial variety (Tjeuw, 2017).

In recent years, works on the genetic improvement of the species have been undertaken but has not yet resulted in improved competitive cultivars (Montes and Melchinger, 2016; Peixoto et al., 2017; Cardoso et al., 2018). For example, in 2018, Cardoso et al. from the *J. curcas* plant breeding program of the Federal University of Viçosa (Brazil) reported the selection of 20 promising individuals for crosses and for cloning aiming to bring genetic gains for successful cultivation. Govender et al. (2018) for their part identified genes with a putative implication in biological processes linked to yield, which could be of great importance in breeding strategies and subsequent production of superior and efficient *J. curcas* varieties.

The present study deals with twenty-two genotypes selected for their production and vigor from a collection of *J. curcas* from Africa, Asia and America, and planted on an experimental farm located in Bokhol (Senegal). For the development of an efficient breeding program, the determination of genetic relationships among these selected genotypes is critical (Díaz et al., 2017). It is an initial step towards efficient parental selection and breeding of superior genotypes. The present study was undertaken to determine and understand the genetic relationships between the selected genotypes, using SSR and AFLP markers in order to propose the best crossing scheme for a breeding program.

## MATERIALS AND METHODS

### Plant materials

The plant material was provided by the Laboratory of Tropical Agroecology of Gembloux Agro-Bio Tech (University of Liège, Belgium). It included dried leaves of twenty-two genotypes of *J. curcas* (Table 1) originating from Africa (Senegal, Burkina Faso, Mali, Congo and Madagascar), Asia (Cambodia, China and India) and America (Ecuador, Dominican Republic and Brazil). These genotypes were selected according to several parameters but especially on the basis of their productivity and their good vigor from a collection of *J. curcas* grown on an experimental farm located in Bokhol in Senegal.

### DNA isolation

Total genomic DNA was isolated from 20 mg of dried leaves following a slightly modified mixed alkyltrimethylammonium bromide (ATAB) method as described by Lacape et al. (2003). Briefly, leaves were ground in 2 ml safe-lock microtubes (Eppendorf, Belgium) with stainless steel beads using a TissueLyser mixer-mill (Qiagen, Belgium) and dissolved in 800 µL of ATAB buffer (0.1 M Tris HCl, pH 8.0, 1.5 M NaCl, 20 mM EDTA, 2% ATAB, 1% polyethylene glycol 6000, 0.5% sodium sulphite) at 72°C. The

**Table 1.** Identification number, country and region of origin of the studied genotypes of *J. curcas*.

N°	Collection identity	Country	Region
1	TA4(13/25)	Senegal	Africa
2	YE4	Senegal	Africa
3	BUR2	Burkina Faso	Africa
4	BA5(1/21)	Burkina Faso	Africa
5	MA3(13/43)	Mali	Africa
6	CMA1	Congo	Africa
7	CMI6	Congo	Africa
8	CMS2	Congo	Africa
9	CMU3	Congo	Africa
10	CON3	Congo	Africa
11	MAD4	Madagascar	Africa
12	MA4	Madagascar	Africa
13	CAM4	Cambodia	Asia
14	CH2	China	Asia
15	IN3	India	Asia
16	INH1	India	Asia
17	INP1	India	Asia
18	MG2	Brazil	America
19	NN2	Brazil	America
20	VE2	Brazil	America
21	EQ2	Ecuador	America
22	RD3	Dominican Republic	America

samples were incubated for 1 h at 72°C and cooled for 5 min at room temperature. Then, 960 µL of chloroform-isoamyl alcohol (CIA) (24:1) was added to each sample. All samples were homogenized by inversion for 5 min, before centrifugation at 6200 g for 20 min at room temperature. The supernatant (800 µL) was collected and the DNA was precipitated with 640 µL of isopropanol by gentle shaking. After a centrifugation at 6200 g for 20 min, the pellet was washed with 600 µL of 70% ethanol through a centrifugation at 6200 g for 10 min, dried at 60°C for 10 min and dissolved in 100 µL of TE buffer (10 mM Tris HCl, pH 8.0, 1 mM EDTA). RNA was removed by RNase treatment at 37°C for 1 h. DNA quality and concentration were evaluated by electrophoresis in SYBR Safe (Invitrogen) stained with 1% agarose gels through comparison with low DNA mass ladder (Invitrogen, Belgium). Final concentration of DNA was adjusted to 50 ng µL<sup>-1</sup> for use in PCR analysis.

### SSR analysis

Ten SSR primer pairs (jcds 10, jcds 24, jcds 58, jcds 66, jcps 1, jcps 6, jcps 20, jcps 21, jcms 2 and jcms 30) were selected from the study of Pamidimarri et al. (2009). Polymerase chain reactions (PCR) were performed in 10 µL volume containing approximately 25 ng of template DNA, 0.6 U of Taq DNA Polymerase, 2.5 mM MgCl<sub>2</sub>, 1x Polymerase Buffer, 2 µM of each forward and reverse primers, and 0.2 mM of dNTPs mix. A PTC-200 thermal cycler (BioRad, Belgium) was used, with a PCR conditions consisting of an initial denaturation at 94°C for 5 min, followed by 35 cycles

of denaturation at 94°C for 30 s, annealing at 55°C for 1 min and extension at 72°C for 1 min, with a final 72°C extension for 8 min. Amplification products were separated on 6.0% denaturing polyacrylamide gel and visualized by silver stain according to the following protocol: Fixing the gel for 3 min in a fixing solution (10% ethanol, 0.5% acetic acid, in water), staining for 5 min in a staining solution (0.2% silver nitrate, in water), rinsing in de-ionized water for 3 s, developing in a cold (4–10°C) developer solution (0.15% sodium hydroxide, 0.2% formaldehyde, in water) until the DNA bands became visible. The gel was rinsed for 1 min in the fixing solution, air-dried and the microsatellite bands were photo-documented and analyzed.

### AFLP analysis

Six AFLP primer combinations were used (E-AAG/M-CAC, E-ACT/M-CAT, E-AAC/M-CTT, E-ACA/M-CAT, E-ACG/M-CTC and E-AGC/M-CAA). AFLP was carried out using the "AFLP Analysis System I / AFLP starter primer kit" (Invitrogen, Belgium) following the protocol proposed by Invitrogen. Briefly, genomic DNA (250 ng) was double digested with *EcoR* I and *Mse* I restriction endonucleases. The digested DNA fragments were ligated to *EcoR* I and *Mse* I adaptors with T4 DNA ligase to generate template DNA for amplification by PCR. Two consecutive PCR were performed: A pre-selective and selective PCR. In the pre-selective reaction, DNA was amplified using an AFLP pre-amp primer pair complementary to the adaptors and each having one selective nucleotide. Pre-selective PCR amplification was used as template for the selective

**Table 2.** Details of the amplified bands obtained from the DNA samples of the twenty two studied genotypes of *J. curcas*, using SSR and AFLP markers.

Marker type	Primer	No. of total bands	No. of polymorphic bands	No. of monomorphic bands	% of polymorphic bands
SSR	jcds 10	8	3	5	37.5
	jcds 24	5	4	1	80
	jcds 58	7	4	3	57.14
	jcds 66	10	4	6	40
	jcps 1	3	0	3	0
	jcps 6	3	1	2	33.33
	jcps 20	9	8	1	88.89
	jcps 21	13	6	7	46.15
	jcms 2	6	4	2	66.67
	jcms 30	11	8	3	72.73
	<i>Total</i>	<i>75</i>	<i>42</i>	<i>33</i>	<i>56</i>
	<b>Mean</b>	<b>7.5</b>	<b>4.2</b>	<b>3.3</b>	<b>56</b>
AFLP	EAAG/MCAC	25	21	4	84
	EACT/MCAT	52	51	1	98.08
	EAAC/MCTT	81	76	5	93.83
	EACA/MCAT	64	62	2	96.87
	EACG/MCTC	53	47	6	88.68
	EAGC/MCAA	93	89	4	95.70
	<i>Total</i>	<i>368</i>	<i>346</i>	<i>22</i>	<i>94.02</i>
	<b>Mean</b>	<b>61.33</b>	<b>57.67</b>	<b>3.67</b>	<b>94.02</b>

amplification using AFLP primers, each containing three selective nucleotides. The amplified fragments were resolved using 6.0% denaturing polyacrylamide gels and DNA bands were visualized by silver staining, as described previously.

#### Data collection and statistical analysis

The scoring of bands was done as present (1) or absent (0) for each SSR and AFLP marker allele and data was entered in a binary data matrix as discrete variables. POPGENE software (version 1.32) was used to calculate observed number of alleles (Na), effective number of alleles (Ne), Nei's genetic diversity (He) and Shannon's information index (I). Within species diversity (Hs), total genetic diversity (Ht), mean coefficient of gene differentiation (Gst) and estimate of gene flow (Nm) were also calculated by POPGENE software. Dissimilarity coefficients between the genotypes, in a pairwise comparison, were computed using the Nei's standard dissimilarity coefficient (Ds) to estimate relationships between the genotypes studied. The resulting dissimilarity matrix was subjected to cluster analysis by Neighbor-joining method and a dendrogram, showing the distance-based interrelationship among the genotypes, was generated using FreeTree software (Pavlicek et al., 1999) and the drawing program TreeView (<http://taxonomy.zoology.gla.ac.uk/rod/treeview.html>).

Analysis of molecular variance (AMOVA) was carried out with GenALEX 6.5 software (Peakall and Smouse, 2001) to examine total genetic variation among and within accessions. In addition, Principal Component Analysis (PCoA) was performed with the

same software, in order to more effectively view the patterns of genetic distance. GenALEX software was also used to calculate Nei's unbiased genetic distance among the different populations. Nei's unbiased genetic distance is an accurate estimate of the number of gene differences per locus when populations are small (Nei, 1978). Finally, a Mantel test was achieved with the GenALEX software to reveal the correlation between the two techniques (SSR and AFLP).

## RESULTS

### Polymorphism of SSR and AFLP amplified products

All the SSR primers and the AFLP primer combinations used amplified fragments across the twenty two genotypes studied (Table 2). In total, 75 bands were revealed for the 10 SSR primers. The number of bands per primer ranged from three (jcps 1 and jcps 6) to thirteen (jcps 21), with an average of 7.5 bands per primer. All the primers showed different levels of polymorphism except jcps 1 which showed no polymorphism among the twenty two jatropha genotypes. The percentage of polymorphic bands per primer ranged from 0% (jcps 1) to 88.89% (jcps 20), with an average of 56%. Four primers out of the 10 primers showed less

**Table 3.** Genetic variability across the three groups of *J. curcas* investigated using SSR primers.

Pop	Observed no. of alleles (Na)	Effective no. of alleles (Ne)	Nei's gene diversity (He)	Shannon's information index (I)
Africa	1.3467 (0.4791)	1.1928 (0.3210)	0.1147 (0.1788)	0.1733 (0.2602)
Asia	1.1200 (0.3271)	1.1031 (0.2919)	0.0541 (0.1510)	0.0766 (0.2123)
America	1.4933 (0.5033)	1.3793 (0.4359)	0.2053 (0.2249)	0.2959 (0.3163)
Africa+Asia+America	1.5600 (0.4997)	1.2347 (0.3060)	0.1490 (0.1709)	0.2354 (0.2503)

The values in the brackets are standard deviation.

**Table 4.** Genetic variability across the three groups of *J. curcas* investigated using AFLP primers.

Pop	Observed no. of alleles (Na)	Effective no. of alleles (Ne)	Nei's gene diversity (He)	Shannon's information index (I)
Africa	1.7962 (0.4034)	1.3503 (0.3571)	0.2122 (0.1816)	0.3318 (0.2475)
Asia	1.4565 (0.4988)	1.2637 (0.3585)	0.1552 (0.1915)	0.2345 (0.2760)
America	1.5924 (0.4921)	1.3162 (0.3530)	0.1910 (0.1872)	0.2924 (0.2688)
Africa+Asia+America	1.9402 (0.2374)	1.3519 (0.3600)	0.2139 (0.1769)	0.3405 (0.2310)

The values in the brackets are standard deviation.

than 50% polymorphic bands. For the six AFLP primer combinations, a total of 368 bands were produced. The number of bands per primer combination ranged from 25 (*EAAG/MCAC*) to 93 (*EAGC/MCAA*) with an average of 61.33 bands per primer combination (Table 2). Of the 368 AFLP bands produced, 346 bands were polymorphic, accounting for 94.02%. The percentage of polymorphism per primer combination ranged from 84% (*E-AAG/MCAC*) to 98.07% (*E-ACT/M-CAT*).

#### Genetic variability revealed with single-population descriptive statistics

At the level of single-population descriptive statistics, SSR results (Tables 3) revealed that the group of American genotypes had the highest values for observed number of alleles (Na), effective number of alleles (Ne), Nei's genetic diversity (He), Shannon's information index (I) with 1.4933, 1.3793, 0.2053 and 0.2959 respectively. The group of Asian genotypes showed the lowest values for these parameters (Na = 1.12, Ne = 1.1031, He = 0.0541 and I = 0.0766), while the group of African genotypes presented intermediate values (Na = 1.3467, Ne = 1.1928, He = 0.1147 and I = 0.1733). The values of these parameters for all the twenty two genotypes were Na = 1.5600, Ne = 1.2347, He = 0.1490 and I = 0.2354.

With AFLP data (Table 4), the highest values for Na, Ne, He and I were found in African genotypes (Na = 1.7962, Ne = 1.3503, He = 0.2122 and I = 0.3318),

followed by American (Na = 1.5924, Ne = 1.3162, He = 0.1910 and I = 0.2924), and then Asian (Na = 1.4565, Ne = 1.2637, He = 0.1552 and I = 0.2345). For all the 22 genotypes the results were Na = 1.9402, Ne = 1.3519, He = 0.2139 and I = 0.3405.

With the combined SSR+AFLP data (Table 5), the highest values for Na, Ne, He and I were found in African genotypes (Na = 1.7201, Ne = 1.3236, He = 0.1957 and I = 0.3050), followed by American (Na = 1.5756, Ne = 1.3269, He = 0.1934 and I = 0.2930), and then Asian (Na = 1.4565, Ne = 1.2637, He = 0.1552 and I = 0.2345). The values of these parameters for the twenty two genotypes were Na = 1.8758, Ne = 1.3321, He = 0.2029 and I = 0.3227.

#### Genetic diversity revealed with multi-populations descriptive statistics

Results of multi-populations descriptive statistics are presented in Table 6. The values for total genetic diversity among the groups (Ht) was 0.1586 for SSR, 0.2095 for AFLP and 0.2009 for SSR+AFLP, while within groups diversity (Hs) was found to be 0.1247, 0.1861 and 0.1757 for SSR, AFLP and SSR+AFLP respectively. Mean coefficient of gene differentiation (Gst) was 0.2152 with SSR data, indicating that 21.52% of the total genetic variation in the groups occurred among groups while 78.48% of the genetic diversity resided within groups. With AFLP and SSR+AFLP data, Gst value was 0.1113 and 0.1252 respectively, indicating that 88.87% (for

**Table 5.** Genetic variability across the three groups of *J. curcas* based on combined SSR+AFLP primers.

Pop	Observed no. of alleles (Na)	Effective no. of alleles (Ne)	Nei's gene diversity (He)	Shannon's information index (I)
Africa	1.7201 (0.4495)	1.3236 (0.3558)	0.1957(0.1846)	0.3050 (0.2564)
Asia	1.3995 (0.4904)	1.2365 (0.3530)	0.1381 (0.1890)	0.2078 (0.2726)
America	1.5756 (0.4948)	1.3269 (0.3685)	0.1934 (0.1939)	0.2930 (0.2770)
Africa+Asia+America	1.8758 (0.3301)	1.3321 (0.3539)	0.2029 (0.1774)	0.3227 (0.2374)

**Table 6.** Values of parameters of multi-populations genetic variability obtained in the study of the three groups of *J. curcas*, based on SSR and AFLP markers.

Marker type	Total gene diversity (Ht)	Intra-population gene diversity (Hs)	Coefficient of gene differentiation (Gst)	Estimate of gene flow (Nm)
SSR	0.1586 (0.0320)	0.1247 (0.0224)	0.2152	1.8234
AFLP	0.2095 (0.0310)	0.1861 (0.0245)	0.1113	3.9920
SSR + AFLP	0.2009 (0.0315)	0.1757 (0.0246)	0.1252	3.4931

The values in the brackets are standard deviation.

**Table 7.** Pairwise population matrix of Nei's unbiased measures of genetic identity (above diagonal) and genetic distance (below diagonal) calculated with combined SSR+AFLP data obtained in the study of the three groups of *J. curcas*.

	Africa	Asia	America
Africa	0.0000	0.9793	0.9763
Asia	0.0209	0.0000	0.9618
America	0.0240	0.0390	0.0000

AFLP) and 87.48% (for SSR+AFLP) of the genetic diversity resided within groups.

Analysis of molecular variance (AMOVA) partitioned the total genetic variance into variance among groups and within groups. The results of AMOVA based on SSR, AFLP and SSR+AFLP data showed respectively 5, 0 and 1% of molecular variance among groups, while within the groups these values were 95, 100 and 99%.

The Nei's unbiased measures of genetic similarity (GS) and genetic distance (GD) among the three studied groups of *J. curcas* were calculated using SSR+AFLP data (Table 7). The GD of the three groups of genotypes varied from 0.0209 to 0.0390, with an average of 0.0279. The largest GD (0.0390) was found between the American and Asian groups and the smallest (0.0209) between African and Asian groups.

#### Genetic relationships among the genotypes and cluster analysis

Genetic relationships among the genotypes of the three

groups of *J. curcas* were constructed by neighbor-joining cluster analysis based on the Nei's standard dissimilarity coefficients (Ds). GD matrixes were obtained from the SSR, AFLP and SSR+AFLP binary data. The Nei's standard dissimilarity coefficients (Ds) between the genotypes ranged from 0.00741 to 0.36682 with a mean of 0.08404 for SSR. The lowest dissimilarity coefficient was between the genotype#10 from Africa (Congo) and the genotype#19 from America (Brazil), and the highest was between genotype#21 from America (Ecuador) and genotype#22 from America (Dominican Republic). For AFLP, the Nei's standard dissimilarity coefficients ranged from 0.22044 to 1.22921 with a mean of 0.53718. The least dissimilar genotypes were genotype#12 from Africa (Madagascar) and genotype#17 from Asia (India) and the most dissimilar were genotype#9 from Africa (Congo) and genotype#11 from Africa (Madagascar). When the SSR and AFLP data were combined, the Nei's standard dissimilarity coefficients ranged from 0.14397 to 0.73943 with a mean of 0.3540. The lowest coefficient was found between genotype#12 from Africa (Madagascar) and genotype#17 from Asia (India), and the highest between

**Table 8.** Genetic distance (GD) matrix obtained by Nei's standard distance (Ds) and neighbor-joining method, showing the relationships among the studied genotypes of *J. curcas* using combined SSR+AFLP data.

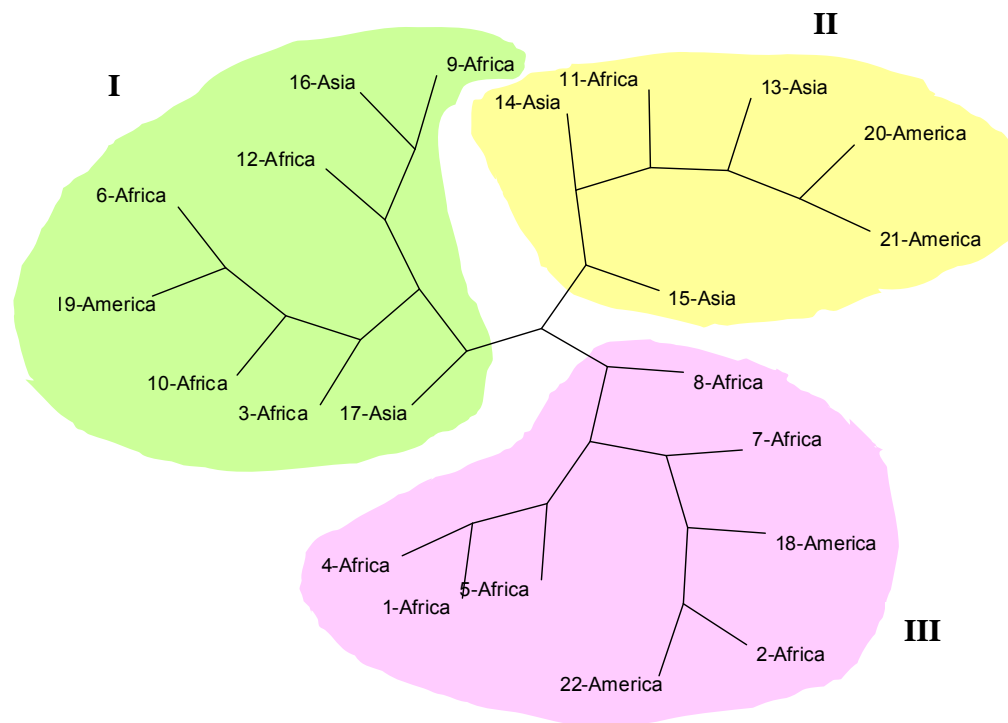
	#1	#2	#3	#4	#5	#6	#7	#8	#9	#10	#11	#12	#13	#14	#15	#16	#17	#18	#19	#20	#21	#22
#1																						
#2	0.229																					
#3	0.354	0.34																				
#4	0.176	0.257	0.334																			
#5	0.326	0.32	0.433	0.374																		
#6	0.327	0.353	0.379	0.371	0.317																	
#7	0.252	0.279	0.391	0.289	0.323	0.326																
#8	0.221	0.243	0.357	0.247	0.311	0.334	0.224															
#9	0.529	0.525	0.561	0.561	0.518	0.5	0.416	0.518														
#10	0.239	0.294	0.282	0.289	0.352	0.319	0.305	0.267	0.549													
#11	0.292	0.307	0.404	0.333	0.414	0.393	0.302	0.258	<b>0.616</b>	0.268												
#12	0.233	0.223	0.317	0.219	0.332	0.353	0.211	0.226	0.459	0.271	0.319											
#13	0.351	0.309	0.396	0.36	0.4	0.403	0.329	0.334	0.515	0.383	0.368	0.312										
#14	0.318	0.288	0.344	0.32	0.325	0.388	0.306	0.283	0.494	0.353	0.363	0.234	0.304									
#15	0.19	0.227	0.353	0.225	0.311	0.318	0.183	0.219	0.475	0.23	0.284	0.17	0.327	0.246								
#16	0.271	0.298	0.411	0.323	0.349	0.323	0.286	0.264	0.456	0.316	0.313	0.259	<b>0.385</b>	0.317	0.262							
#17	0.193	0.211	0.336	0.243	0.313	0.341	0.199	0.2	0.5	0.251	0.255	0.144	0.273	0.267	0.158	0.262						
#18	0.268	0.286	0.35	0.297	0.339	0.306	0.19	0.268	0.391	0.26	0.341	0.226	0.334	0.276	0.206	0.346	0.221					
#19	0.31	0.266	0.349	0.358	0.321	0.352	0.294	0.341	0.482	0.31	0.412	0.267	0.363	0.361	0.271	0.382	0.223	0.308				
#20	0.446	0.451	0.514	0.504	0.435	0.491	0.39	0.43	0.615	0.463	0.461	0.372	0.451	0.425	0.388	0.419	0.434	0.421	0.548			
#21	0.421	0.453	0.522	0.478	0.454	0.463	0.456	0.395	<b>0.739</b>	0.421	0.465	0.438	0.525	0.473	0.398	0.45	0.452	0.461	0.58	0.518		
#22	0.394	0.363	0.504	0.425	0.407	0.421	0.384	0.416	0.546	0.375	0.501	0.33	0.434	0.392	0.361	0.368	0.355	0.317	0.374	0.433	<b>0.604</b>	

genotype#9 from Africa (Congo) and genotype#21 from America (Ecuador). The GD matrixes generated by the combined SSR+AFLP data (Table 8) also showed that, the most genetically distant genotypes in the African group were genotype#9 from Congo and genotype#11 from Madagascar (Ds = 0.61553). In the American groups the most distant genotypes were genotype#21 from Ecuador and genotype#22

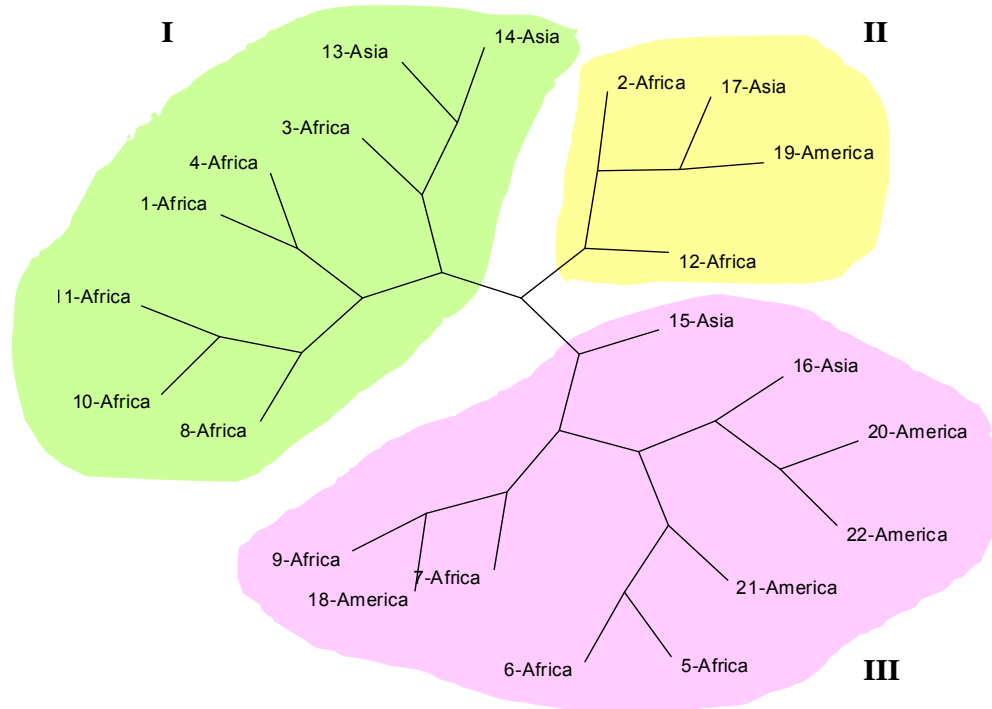
from Dominican Republic (Ds = 0.60396). And finally in the Asian group, the most distant genotypes were genotype#13 from Cambodia and genotype#16 from India (Ds = 0.38502).

The dendrograms (Figures 1 to 3) constructed, defining the genomic relationships among analyzed genotypes, grouped the 22 *J. curcas* genotypes into three main clusters (I, II and III). For SSR, the composition of each cluster (Figure

1) was: Eight genotypes for cluster I (2 from Asia, 1 from America and 5 from Africa), six genotypes for cluster II (3 from Asia, two from America and 1 from Africa) and eight genotypes for cluster III (2 from America and 6 from Africa). The clustering results for AFLP (Figure 2) was: Eight genotypes for cluster I (2 from Asia and 6 from Africa), four genotypes for cluster II (1 from Asia, 1 from America and 2 from Africa), ten genotypes for

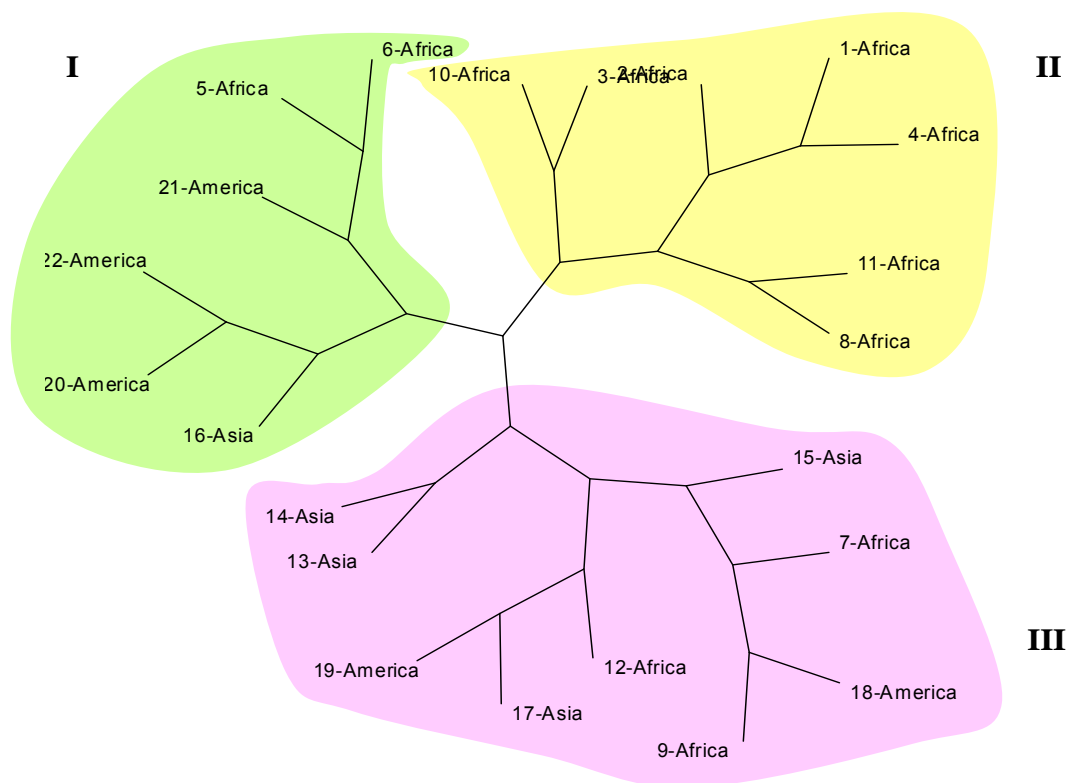


**Figure 1.** Dendrogram generated using Nei's standard distance (Ds) and neighbor-joining method, showing relationships between studied genotypes of *J. curcas*, based on SSR markers.



**Figure 2.** Dendrogram generated using Nei's standard distance (Ds) and neighbor-joining method, showing relationships between studied genotypes of *J. curcas*, based on AFLP markers.





**Figure 3.** Dendrogram generated using Nei's standard distance (Ds) and neighbor-joining method, showing relationships between studied genotypes of *J. curcas*, based on AFLP+SSR markers.

cluster III (2 from Asia, 4 from America and 4 from Africa). For combined SSR and AFLP data, the clustering (Figure 3) was: Six genotypes for cluster I (2 from Africa, 1 from Asia and 3 from America), seven genotypes from Africa for cluster II and nine genotypes for cluster III (3 from Africa, 4 from Asia and 2 from America).

Principal Coordinate Analysis (PCoA) was used to add complementary information to the cluster analysis. The results of PCoA (Figure 4a, b and c) showed, clustering comparable to those of the dendrograms. No clear relation was shown between genotype associations and geographical origin.

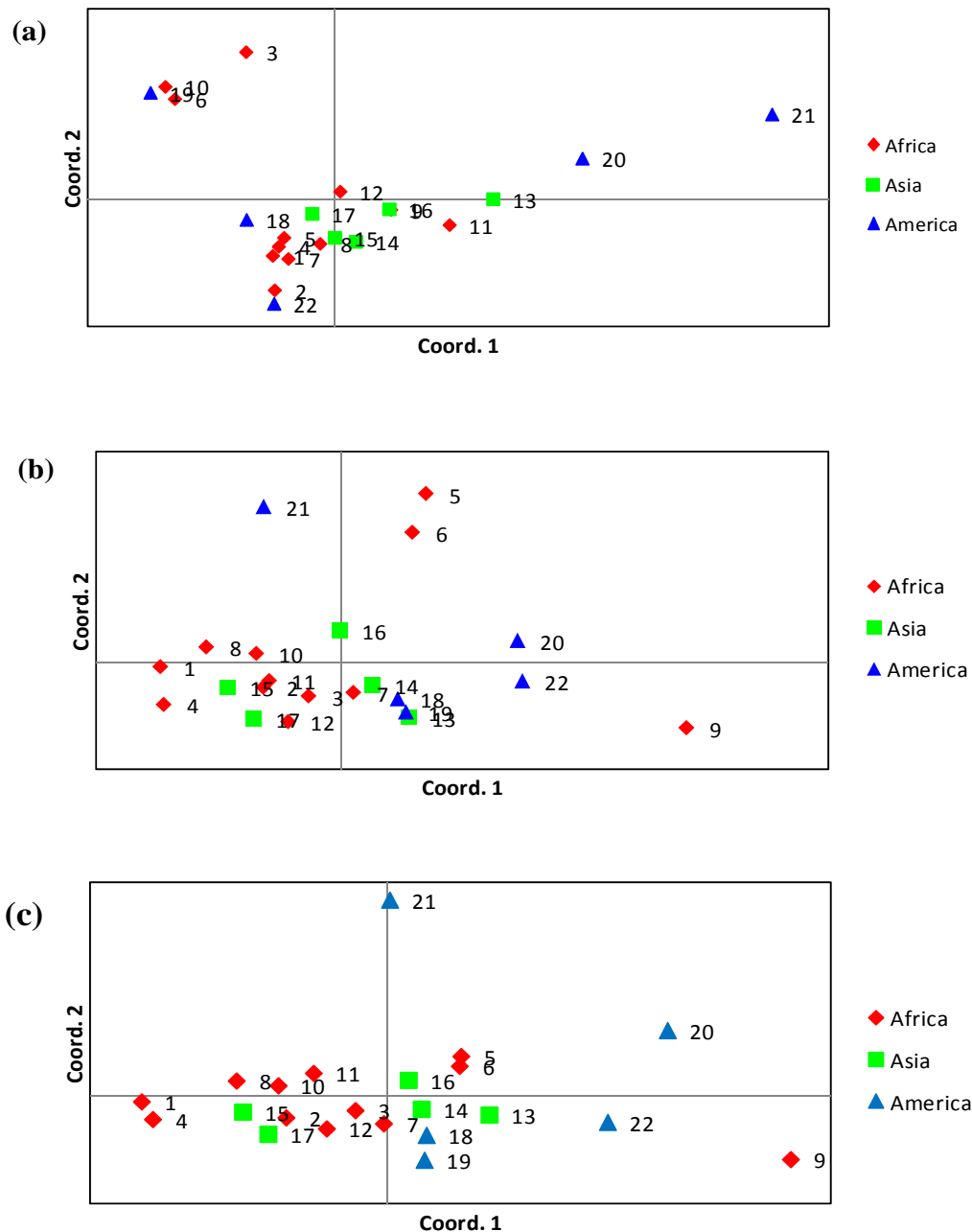
A mantel test between the two Nei's genetic diversity indexes was performed with 99 permutations, in order to estimate correlations for the two molecular systems. It resulted in  $r^2 = 0.1093$  (Figure 5).

## DISCUSSION

In the present study, the two marker approaches (SSR and AFLP) showed their ability to effectively reveal polymorphism among plant materials of *J. curcas*. However, AFLP exceeded SSR in the ability to detect

genetic polymorphism with higher resolving power (94.02% of polymorphic bands for AFLP against 56% for SSR). This might be due to the type of genetic polymorphism detected by each molecular marker system and the different type of information provided by each of them (Augustinos et al., 2016). Indeed, the higher multiplexing ability of arbitrarily dominant technologies such as AFLP is more efficient in detecting polymorphism per assay rather than high levels of polymorphism at each locus of the codominant SSR technology (Costa et al., 2016). Osorio et al. (2014) using a set of 29 SSR and 20 AFLP in a collection of *J. curcas*, found also higher percentage of polymorphism with AFLP (86%) than with SSR (73%).

The difference between the two marker systems was also shown by the Mantel test that revealed no significant associations between them ( $r^2 = 0.1093$ ). A low correlation between AFLP and SSR has also been reported by Mardi et al. (2011) working on durum wheat, with  $r^2 = 0.116$ . According to Augustinos et al. (2016), poor correlation between genetic markers most likely indicates that they refer to different portion of the genome. Accordingly, the combined use of different marker systems may provide more reliable information

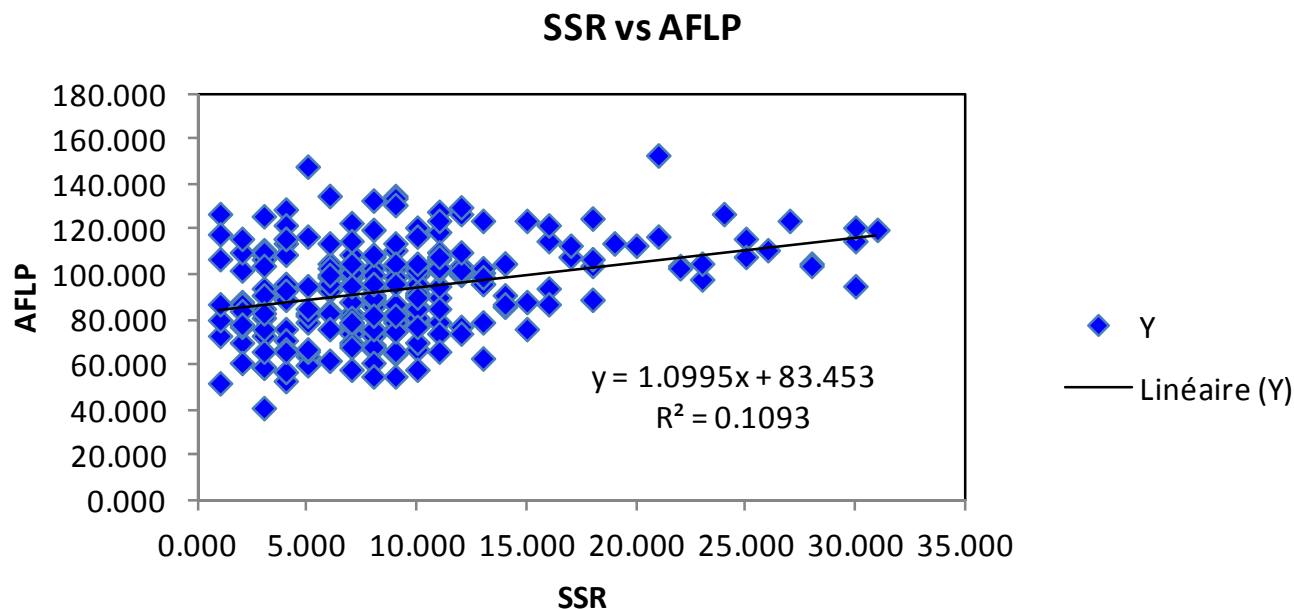


**Figure 4.** Two-dimensional plot of principal coordinate analysis (PCoA) of the twenty two studied genotypes of *J. curcas*, using SSR (a), AFLP (b) and combined SSR+AFLP (c) data. The numbers plotted represent genotype N°.

about genetic diversity when compared to the use of only one marker system (Leal et al., 2010; Augustinos et al., 2016). Therefore, the results obtained from the two marker systems used in the present study should be considered complementary.

With the combined data of the two markers systems, the values of Nei's genetic diversity obtained for the

twenty two selected genotypes was  $H_e = 0.2029$ . This value was lower than those reported from populations of *J. curcas* in Chiapas in Mexico ( $H_e = 0.34-0.54$ ) by Sanou et al. (2015). Considering that Chiapas (Mexico) is recognized as a center of origin and diversity of *Jatropha* (Angel et al., 2016), the genetic diversity of the selected genotypes in the present study was not negligible.



**Figure 5.** Output of mantel test comparing the AFLP and SSR genetic distance matrices.

Wright (1951) has previously shown that  $G_{st}$  is a valuable parameter to determine the degree of genetic differentiation between populations. For the selected genotypes,  $G_{st}$  value was 0.1252 with combined SSR+AFLP. This  $G_{st}$  value globally indicated that more than 87.48% of the genetic diversity resided within the groups and only 12.52% of the total genetic variation occurred among groups. These results agreed with the weak molecular variance among groups shown by AMOVA analysis and the low genetic distances found among the three groups of genotypes. Other studies in *J. curcas* based on different types of molecular markers have also reported that in general, there is limited variation among groups of *J. curcas* from various part of the world (Kumar et al., 2009; Subramanyam et al., 2009; Rosado et al., 2010). The low genetic differentiation among groups indicates that accessions of *J. curcas* have originated from the same genetic background, as mentioned by Pamidimarri and Reddy (2014) and Díaz et al. (2017).

Analysis of the genetic relationships between genotypes using the Nei standard dissimilarity matrix obtained from combined SSR + AFLP data, showed dissimilarity coefficients ranging from 0.14397 to 0.73943. This result indicated the existence of relatively important genetic distances between some genotypes. The most distant genotypes can be good candidates for crosses. The dendrogram, obtained from these genetic relationships among the genotypes, grouped the 22 *J. curcas* genotypes into three main clusters (I, II and III).

Globally, the dendrogram showed three tendencies: (i) Genotypes from diverse regions group together in the same cluster, (ii) Genotypes from the same region scattered in different clusters, and (iii) Genotypes from the same region cluster together. This pattern of clustering showed no clear association between geographical distribution of genotypes and genetic diversity. This means that the geographical origin of the genotypes studied is not significantly related to their genetic similarity. Principal Coordinate Analysis (PCoA) confirmed this clustering pattern. Similar results were found by other authors such as Ambrosi et al. (2010), Maghuly et al. (2015) and Konan et al. (2018) showing that geographical origin do not necessarily reflect the genetic diversity of accessions of *J. curcas*. This lack of correspondence between the molecular classification and geographic origin of the accessions could be due to their possible common origin, confirming the hypothesis that *J. curcas* genetic stocks of Asia and African were introduced from America (Li et al., 2017; Gangapur et al., 2018).

The challenge of all plant genetic breeding programs is to maintain the genetic diversity within the target species while improving desired traits that enable plant materials to perform well (Díaz et al., 2017). The results obtained in this study indicate the existence of a certain level of diversity that can be used by breeders. The choice of the parents can be based on the distance existing between the different genotypes. Based on the Nei's standard dissimilarity matrix and the PCoA analysis using the

combined SSR+AFLP data, the greatest heterosis might be expected from the following crossing schemes. Cross on the one hand, the two most genetically distant genotypes from Africa (genotype#9 from Congo and genotype#11 from Madagascar); and on the other hand, cross the two most genetically distant genotypes from America (genotype#21 from Ecuador and genotype#22 from Dominican Republic). Then, select from each cross, the best F1 progenies by phenotypic performance. Finally, cross the selected African F1 genotypes to the selected American F1 genotypes and select their best progenies according to their phenotypic performances. These selected progenies can be multiplied by vegetative propagation to preserve excellent clonal genetic stocks in the heterozygous status or used for further selections by back crossing. Such crossings might allow for greater success in the production of genetic variability and thus might maximize the exploitation of heterosis and segregation.

## CONFLICT OF INTERESTS

The authors have not declared any conflict of interests.

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